ABSTRACT: In nature, co-assembly of polypeptides, nucleic acids, and polysaccharides is used to create functional supramolecular structures. Here, we show that DNA nanostructures can be used to template interactions between peptides and to enable the quantification of multivalent interactions that would otherwise not be observable. Our functional building blocks are peptide–oligonucleotide conjugates comprising de novo designed dimeric coiled-coil peptides covalently linked to oligonucleotide tags. These conjugates are incorporated in megadalton DNA origami nanostructures and direct nanostructure association through peptide–peptide interactions. Free and bound nanostructures can be counted directly from electron micrographs, allowing estimation of the dissociation constants of the peptides linking them. Results for a single peptide–peptide interaction are consistent with the measured solution-phase free energy; DNA nanostructures displaying multiple peptides allow the effects of polyvalency to be probed. This use of DNA nanostructures as identifiers allows the binding strengths of homo- and heterodimeric peptide combinations to be measured in a single experiment and gives access to dissociation constants that are too low to be quantified by conventional techniques. The work also demonstrates that hybrid biomolecules can be programmed to achieve spatial organization of complex synthetic biomolecular assemblies.

KEYWORDS: self-assembly, DNA nanostructure, peptide, heterodimeric coiled coil, polyvalent binding, peptide–oligonucleotide conjugate

Natural biological molecules—lipids, nucleic acids, peptides, proteins, and polysaccharides—fold, assemble, and co-assemble to form a stunning variety of functional three-dimensional structures and complexes. Understanding and harnessing these processes underpins emerging applications in synthetic biology and biotechnology.1–3 Broadly speaking, there are two approaches to the manipulation of biomolecular assembly. Biomolecular engineering allows the chemistries of natural biomolecules to be altered to produce variants of wild-type parents whose interactions are tailored to specific applications.4–6 The alternative, de novo design, is completely bottom-up: polypeptide or nucleic acid sequences are generated from scratch by the designer to fold and assemble in prescribed ways.2,3,7,8 This second route has the potential to access a much larger design space but requires clear and reliable sequence-to-structure design rules: this is particularly challenging in the case of polypeptides. To begin to approach the complexity of natural systems and to harness fully the potential of de novo biomolecular assembly, we must learn to control and combine the assembly of different biomolecular types to generate structured, functional co-assemblies.9,10

The specificity of Watson–Crick pairing between the four DNA bases has enabled the fabrication of a rapidly increasing range of DNA nanostructures, especially DNA origami nanostructures, whose assembled shapes can be specified with high precision and reliability through design of the oligonucleotide components.11–18 This process is facilitated by rapid access to synthetic oligonucleotides with ever-decreasing price,19 the application of computer-aided design,20,21 and,

**Supporting Information**

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most recently, the scaling up of DNA production and assembly. However, although well adapted to programmed nanostructure assembly, DNA has minimal intrinsic functionality. In contrast, the chemical diversity of the natural amino acids underpins the folding of a wide variety of protein structures with highly evolved functions including architecture, signaling, active and passive transport, and catalysis. However, the relationship between the amino acid sequence of a polypeptide and its structure and function is complex, which makes de novo protein design extremely challenging.

For certain classes of protein structure, enough is understood to allow rational and reliable de novo design. For example, for α-helical coiled coils, which are bundles of two or more α-helices, established sequence-to-structure relationships have enabled the design of homomeric dimers, trimers and tetramers and the assembly of obligate heterodimers and trimers. More complex and elaborate architectures—for instance, antiparallel arrangements of helices and homomeric and heteromeric assemblies above a tetramer—have proved more challenging to design and have been discovered serendipitously or designed computationally.

Scheme 1. Assembly of Megadalton DNA Origami Structures Is Driven by Peptide–Oligonucleotide Conjugates through Formation of Peptide Coiled-Coil Heterodimers and Pairing of Oligonucleotide Tags with Complementary ssDNA Handles

Peptide–oligonucleotide conjugates are formed via a Cu-free cycloaddition between the highlighted azide side chains of the peptides and the ring-strained dibenzylcyclooctyne groups of the oligonucleotide tags.
way to access more complex structures is to make use of the high spatial resolution and predictable architectures of DNA nanostructures to template peptide assemblies and thus force otherwise unobtainable configurations of α-helices. Others have described the assembly of trimeric coiled-coil DNA conjugates, layered nanostructures formed from collagen-mimetic peptides and oligonucleotides, and DNA-directed peptide pores; conversely, proteins have been used to direct the folding of DNA scaffolds.

Here, we describe the formation of peptide–oligonucleotide conjugates that can be used to control the assembly of larger DNA–peptide structures and thereby to quantify multivalent peptide–peptide interactions. Single-stranded DNA (ssDNA) “handles”, extending from two distinct DNA origami nanostructures, hybridize to complementary oligonucleotide “tags” which are covalently linked to peptides. This places a prescribed number of peptides, each of which is half of a heterodimeric peptide coiled coil, on one end of each DNA origami. Dimerization of the composite nanostructures is driven by association of the peptides to form coiled coils (Scheme 1). Markers on the DNA origamis allow assembled species to be distinguished by transmission electron microscopy (TEM). Counting the distributions of all kinds of assembly in electron micrographs allows the dissociation constant ($K_d$) for each to be estimated and the effect of polyvalency on peptide–peptide interactions to be investigated.

RESULTS AND DISCUSSION

Peptides Compatible with Both Oligonucleotide Conjugation and Peptide Assembly Can Be Designed De Novo. Coiled-coil peptide sequences have heptad repeats, with hydrophobic residues predominating at the “a” and “d” sites. These form amphipathic α-helices whose assembly is driven by the formation of a d/h hydrophobic seam. Judicious placement of specific amino acids at these positions dictates an oligomeric state, with a = isoleucine (Ile) and d = leucine (Leu) directing dimers. The “e” and “g” sites which flank the hydrophobic core can be used to stabilize both homo- and heteromeric assemblies.

Complementary patterns of charged residues at these sites have been used by many groups to create de novo heterodimers.
The simplest previously reported coiled-coil heterodimers have glutamic acid (Glu) at all “e” and “g” sites in one peptide and lysine (Lys) at these sites in the complementary peptide. However, we confirmed that the basic peptide binds nonspecifically and tightly to DNA, consistent with strong electrostatic interactions, making it unsuitable for use in composite peptide–DNA nanostructures. We therefore designed a series of parallel heterodimeric coiled coils in which each peptide was, overall, close to or exactly charge-neutral at pH 7. Peptide sequences are given in Supporting Information (SI), Table S1. These peptides had blocks of both Glu- and Lys-based heptads, with Asn residues at some of the central “a” sites to add stability and increase specificity. They varied in length, having 3, 3.5, or 4 heptad repeats and, in the case of the 4-heptad pairs, either one or two asparagine residues at “a” sites to destabilize off-target homomers. The peptides were made by solid-phase peptide synthesis, purified by high-performance liquid chromatography (HPLC), and confirmed by matrix-assisted laser desorption/ionization time of flight mass spectrometry (SI Figure S1). They were characterized in solution, both alone and in combination with their designed partners, using circular dichroism (CD) spectroscopy and analytical ultracentrifugation (AUC) (Figure 1 and SI Figures S2 and S3, Tables S2 and S3).

On the basis of CD and AUC characterization, the CC-Di-EK and CC-Di-KE pair (Figure 1), comprising four complete heptad repeats with asparagines at two “a” sites, was identified as a suitable coiled coil to link peptide–DNA nanostructures. CD spectra showed that unmixed CC-Di-EK and CC-Di-KE were either largely or fully unfolded at 5 °C, neutral pH, and 50 μM concentration, but that a 1:1 mixture of the peptides displayed α-helical and had a sigmoidal thermal unfolding transition with a midpoint (T_m) of 43 °C (Figure 1a,b), consistent with the formation of a stable coiled-coil heterodimer. From the concentration dependence of the T_m the K_d for heterodimerization was estimated to be 102 ± 26 nM (SI Figure S4). Fits of AUC data assuming a single ideal species returned the mass of the CC-Di-EK:KE dimer (Figure 1c) and confirmed that the unmixed peptides were monomers (SI Figure S3 and Table S3). Molecular models built and optimized in ISAMBARD indicate that these peptides form a stable coiled-coil dimer with a strong preference for the parallel helix orientation (Figure 1d and SI Figure S5). To confirm the preferred helix orientation, the peptides were synthesized with terminal Cys residues and linked as obligate heterodimers via disulfide bonds. With the peptides linked via the same ends—that is, both C-termini, consistent with parallel helix formation—the resulting conjugate was highly helical, thermally stable, and monomeric (SI Figures S2 and S3, Tables S2 and S3). In contrast, when linked via opposite ends, the behavior of the construct was more complex with larger oligomeric complexes indicated by AUC. These experiments support the conclusion that the unlinked CC-Di-EK and CC-Di-KE peptides form a stable, parallel, helical heterodimer. CD spectra showed no interaction between either CC-Di-EK or CC-Di-KE and the ssDNA tags (SI Figure S6a,b). This confirmed our design choice to use charge-neutral or acidic peptides for conjugation to DNA tags.

Other peptide pairs were not developed further. CD and AUC experiments (SI Figures S2 and S3, Tables S2 and S3) showed that the 4-heptad peptides with only a single asparagine displayed off-target homomeric interactions and that the 3-heptad pair was not fully folded. The 3.5-heptad pair CC-Di-EN3.5 was rejected because it had a +1 net charge. Peptide–oligonucleotide conjugates were synthesized as follows. Azidonorleucine (Z) was added during peptide synthesis at the C-terminus of CC-Di-EK and at the N-terminus of CC-Di-KE to give CC-Di-EK-Z and Z-CC-Di-KE (SI Table S1). These were then conjugated to dibenzylcyclooctyne-functionalized oligonucleotide tags, α and β, respectively, by copper-free 1–3 Huisgen cycloaddition (Scheme 1 and SI Figure S6). The resulting conjugates,
CC-Di-EK-α and β-Cc-Di-KE, were isolated by reverse-phase HPLC and confirmed by LC-MS (SI Figures S6 and S7). Flexible single-stranded spacers (eight thymidines) were included in the oligonucleotide tags to separate the peptide attachment site and the duplex formed between tag and complementary handle (SI Figure S10 and Table S6). The CD spectra of the peptide−oligonucleotide conjugates when mixed were consistent with the formation of a fully folded α-helical species and had a thermal unfolding profile ($T_M = 34 \, ^\circ C$) similar to that of the untagged peptides (SI Figure S8: the predicted $T_M$ for unconjugated peptides at this concentration is $36 \, ^\circ C$).

Two Distinct DNA Nanostructures Whose Identities and Orientation Can Be Determined by Electron Microscopy. Two readily distinguishable, asymmetric DNA origami nanostructures, A and B, were designed with the assistance of caDNAno software.20,21,46 (Figure 2a,b and SI Figure S9). Each had a central six-helix bundle with either one or two asymmetrically placed sleeves (bulges comprising a concentric layer of DNA helices) wrapped around it. The nanostructures were assembled by annealing a 2686 nucleotide single-stranded scaffold, derived from pUC19 plasmid,48 with 69 and 67 synthetic oligonucleotide staples, respectively (SI Table S6). The asymmetric positions of the sleeves allowed rapid determination of the identity and orientation of individual origamis in electron micrographs (Figure 2c,d).

Each origami was functionalized at the end further from the sleeves with three single-stranded 23 nucleotide handles as extensions to selected staples (Figure 2a,b). Extensions were either complementary to the oligonucleotide tags α and β, referred to as A and B handles, respectively, or non-complementary dummy sequences (SI Figure S10 and Table S6).

Peptide−Oligonucleotide Conjugates Bring Together DNA Nanostructures. Each DNA origami was decorated with one, two, or three copies of the corresponding half of the heterodimeric coiled-coil peptide pair, that is, A with CC-Di-EK-α and B with β-Cc-Di-KE. Preassembled origamis

Figure 3. Directed assembly of DNA nanostructures through interactions of peptide−oligonucleotide conjugates. (a) DNA origamis A, and B, decorated with three peptides. (b–e) Representative TEM images of the assembly of peptide-decorated DNA origamis. The distribution between species depends on the number of peptides, $n$, displayed on each origami: (b) $n = 3$, (c) $n = 2$, (d) $n = 1$, and (e) $n = 0$. White numbers in (b) indicate representative heterodimers (1 and 2) and monomers (3 and 4). The initial concentrations of assemblies A and B were, respectively, 0.61 and 0.84 nM (b, $n = 3$), 0.84 and 1.09 nM (c, $n = 2$), 1.69 and 2.39 nM (d, $n = 1$), and 0.68 and 0.74 nM (e, $n = 0$). Scale bars: 50 nm. (f) Dissociation constants of heterodimers of origamis functionalized with different numbers of peptides were determined from linear fits (solid lines) to plots of $[A_n] \times [B_n]$ as a function of $[A_n]$, over a range of total monomer concentrations from 0.25 to 2.4 nM. Data were weighted by estimated uncertainties (SI Tables S4, S5 and supporting discussion).
displaying the specified number of complementary ssDNA handles were incubated at room temperature (approximately 20 °C) with a 3× stoichiometric excess of the corresponding oligonucleotide-tagged peptide. The resulting peptide-decorated origamis, named \( A_n \) and \( B_n \) where \( n (n = 1, 2, 3) \) indicates the number of peptides attached to each DNA origami, were purified by gel electrophoresis (Figure 3a and SI Figure S11). Subsequent mixing of purified \( A_n \) and \( B_n \) resulted in the formation of heterodimeric complexes \( A_n:B_n \) which, when imaged by TEM, were seen to be joined almost exclusively via their peptide-labeled ends (Figure 3b–d and SI Figure S12), consistent with dimerization driven by the formation of peptide coiled coiled as designed.

The use of two DNA origami structures that can be distinguished in electron micrographs allowed us to identify manually and count free assemblies \( A_n \) \( B_n \) and all complexes formed between them, that is, the intended heterodimer \( A_n:B_n \), homodimers \( (A_n:A_n \) and \( B_n:B_n \), and any larger multimers (Figure 3b–e and SI Figure S12 and Table S4). If it is assumed that the distribution between complexes on the TEM grid is representative of that present in solution, then, given the initial total concentrations of the two peptide-decorated DNA origamis, the equilibrium concentration of each species can be inferred. In each experiment, approximately 1000 origamis of each type were counted and classified as monomeric, homor heterodimeric, or as part of a larger multimer. Uncertainties in inferring equilibrium concentrations of these species from the observed distributions of particles in electron micrographs are discussed in the SI supporting discussion and quantified in SI Table S5.

As anticipated, in the absence of attached peptides \( (n = 0) \), very few complexes were counted (Figure 3e and SI Figure S12-1 and Table S4-1). This background count is likely to be due to chance observations of two origami structures in close proximity rather than to any interaction between the origamis: the data are consistent with a random distribution of origami nanostructures in each micrograph with a minimum detectable separation between the ends of adjacent origamis of 8 nm (SI supporting discussion). This minimum radius was used to calculate a background due to random coincidences which was subtracted from the observed numbers of dimers between nanostructures functionalized with one, two, or three peptides (SI Tables S4-2, S4-3, and S4-4).

For mono- and bifunctionalized origamis \( (n = 1, 2) \), the numbers of dimers observed were small, typically <2 and <4%, respectively, of the number of monomers counted, but for trifunctionalized origamis, the population of heterodimers reached 40% of the total number of particles. It was possible to obtain consistent estimates of the dissociation constants for all heterodimers and for \( n = 3 \) homodimers. \( K_d \) values were estimated from the fitted slopes of plots of [\( A_n \) \( B_n \)] vs [\( A_n:B_n \), [\( A_n \)] vs [\( A_n:A_n \), and [\( B_n \)] vs [\( B_n:B_n \)] for a range of initial monomer concentrations (0.25–2.4 nM) (Figure 3f and SI Figure S13). Results are presented in Table 1.

Our estimated \( K_d \) of 130 ± 20 nM for the monofunctionalized heterodimer \( A_1:B_1 \) is close to the value of 102 ± 26 nM determined from the melting temperatures of free peptides in solution (SI Figure S4). This is consistent with our design hypothesis that interactions between peptide-functionalized origamis are dominated by specific peptide–peptide interactions and suggests that the preparation of samples for electron microscopy does not introduce significant sampling bias (SI supporting discussion). Increasing the number of displayed peptides increases the interaction strength, decreasing the dissociation constant by approximately 200× for the trifunctionalized heterodimer. Association between trifunctionalized homodimers \( A_3:A_3 \) and \( B_3:B_3 \) is clearly resolved and is 30× weaker than for the corresponding heterodimer, consistent with the designed association bias of the peptides.

Upon increasing the number of attached peptides from \( n = 1 \) to \( n = 2 \), the heterodimer \( K_d \) decreases only by a factor of approximately 5, which is orders of magnitude less than the factor that would be expected if two parallel peptide links could be formed independently.\(^{49,50}\) As shown in SI Figures S9 and S10, the ends of the origamis are not flat (the ends of the six bundled DNA helices are not aligned). Despite the presence of flexible T\(_4\) linkers between each peptide and the DNA duplex linking it to the origami, this may hinder the simultaneous formation of multiple coiled-coil links between pairs of origamis, providing a natural explanation for the unexpectedly small increase in affinity observed. The observed increase in affinity can be almost entirely attributed to the 4-fold increase in the number of ways of making a single coiled coil from two pairs of peptides, consistent with the hypothesis that a single peptide–peptide link dominates interactions between doubly functionalized origamis. If this combinatorial argument were extended to the \( n = 3 \) system, we would predict \( K_d \approx 10\) nM, as there are nine ways to make a single coiled-coil link between origamis displaying three peptides each. However, we observed an order of magnitude greater enhancement. This strongly suggests polyvalent binding:\(^{49,50}\) there is a significant probability that at least two of the three possible coiled-coil links are formed simultaneously.

**CONCLUSIONS**

In summary, we have shown that DNA origami nanostructures can be used to template and quantify the assembly of de novo designed peptides. To avoid nonspecific binding of the peptides to DNA, it was necessary to design charge-neutral, heterodimeric coiled-coil peptides which will prove useful in biomolecular construction and synthetic biology more generally. The use of distinguishable DNA origami conjugates allows the use of electron microscopy to estimate otherwise undetectable concentrations of free and bound peptides. The direct enumeration of interacting species provides an alternative way to determine \( K_d \) which can be applied at concentrations much lower than would be practical for traditional methods such as CD spectroscopy or isothermal calorimetry. It also allows simultaneous measurement of dissociation constants for all observed species in a reaction mixture, in this case, homomeric and heteromeric complexes.

**Table 1. Dissociation Constants of Peptide-Mediated DNA Origami Dimers Inferred from the Relative Numbers of Complexes Observed in Electron Micrographs as Functions of Total Monomer Concentrations**

<table>
<thead>
<tr>
<th>Valency, ( n )</th>
<th>Complex ( A_n:B_n )</th>
<th>Complex ( A_n:A_n )</th>
<th>Complex ( B_n:B_n )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>130 ± 20</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>25 ± 3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>0.7 ± 0.1</td>
<td>24 ± 4</td>
<td>20 ± 3</td>
</tr>
</tbody>
</table>

\(^{49,50}\) See SI supporting discussion of experimental errors: there is an additional systematic error of approximately 20% in all \( K_d \) values corresponding to uncertainty in determining DNA origami concentrations.
something that would be impossible to deduce by measurement of a single average quantity such as circular dichroism. This semiquantitative technique also allows us to probe the effects of polyclonal interactions: the enhanced binding of trisfunctionalized origami heterodimers indicates that the use of a DNA origami template to preposition multiple peptides is an effective method to control their interactions. This study shows the way to the development of more sophisticated DNA templates to control the spatial orientation of peptide and protein assemblies and to the use of peptide–peptide interactions to direct DNA assembly.

EXPERIMENTAL SECTION

Peptide Synthesis. Peptides were synthesized via solid-phase peptide synthesis on a rink amide resin (Novabiochem, 0.1 mmol scale) using a microwave-assisted CEM Liberty Blue synthesizer (CEM Corporation) and standard deprotection and coupling chemistries. Peptide-grade dimethylformamide (DMF) was used throughout. Fmoc deprotections were performed with 20% morpholine in DMF (Merck-Millipore), followed by CI-HOBt (AGTC Bioproducts, 0.5 M in DMF) and DIC (AGTC Bioproducts, 1 M in DMF) activated couplings of each amino acid (AGTC Bioproducts). N-Terminal acetylations were performed with pyridine (Fisher Scientific, 0.5 mL) and acetic anhydride (BDH Laboratories, 0.25 mL) in DMF (15 min). Peptides were cleaved from the resin in trifluoroacetic acid (Acros Organics) with a trisopropylsilane scavenger (Sigma-Aldrich) and H₂O at 90:5:5 vol % under agitation for 2 h. The cleavage mixtures were then filtered to remove the resin, and the volume was reduced to <5 mL by a flow of nitrogen before precipitation of the crude peptides by addition of cold diethyl ether. A solid peptide pellet was obtained by centrifugation followed by removal of the supernatant. This was then lyophilized from 1:1 H₂O/MeCN to yield crude peptides as white powders.

DNA Origami Design. DNA origamis A and B were designed using cadDNAo. Nicks separating staple strands were positioned manually, as shown in SI Figure S9. As shown in SI Figure S10, staple strands on helices 0, 2, and 4, at the end further from the marker N-Terminal acetylations were performed with pyridine (Fisher Scientific) and DIC (AGTC Bioproducts) following by CI-HOBt (AGTC Bioproducts). N-Terminal acetylations were performed with pyridine (Fisher Scientific) and acetic anhydride (BDH Laboratories) in DMF. Peptides were cleaved from the resin in trifluoroacetic acid (Acros Organics) with a trisopropylsilane scavenger (Sigma-Aldrich) and H₂O at 90:5:5 vol % under agitation for 2 h. The cleavage mixtures were then filtered to remove the resin, and the volume was reduced to <5 mL by a flow of nitrogen before precipitation of the crude peptides by addition of cold diethyl ether. A solid peptide pellet was obtained by centrifugation followed by removal of the supernatant. This was then lyophilized from 1:1 H₂O/MeCN to yield crude peptides as white powders.

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Equilibrium Concentration Calculations. The numbers (N) of monomers (A₁, B₁) and of dimers (A₂, B₂, A₁B₂, B₁A₂) and higher multimers, in which the proximity of the peptide-functionalized ends was consistent with the formation of a peptide-bound complex, were counted. A background corresponding to the random chance of observing two origami structures in close proximity (i.e., when their peptide-functionalized ends are within 8 nm of each other; see main text) was subtracted. The equilibrium concentration of each species was assumed to be proportional to the number counted and was estimated from counted numbers of each assembled complex and the total monomer concentrations (summed over all assemblies) (SI Table S5). Kd Calculation and Fitting Model. Linear fits to graphs of [A₁] × [B₁] vs [A₁B₂], [A₁]² vs [A₂], and [B₁]² vs [B₁A₂] for six different initial concentrations of [B₁] × [B₁] from 0.25 to 2.4 nm were obtained using OriginPro (Figure 3f and SI Figure S13). Data were weighted by estimated uncertainties in both quantities (SI supporting discussion and Table S5). The dissociation constant was estimated as the gradient of a linear fit to these data which was constrained to pass through the origin, (0,0).

ASSOCIATED CONTENT

Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsnano.9b04251.

Additional methods; peptide interaction models; synthesis of peptide–oligonucleotide conjugates; biophysical characterization of peptides and peptide–oligonucleotide conjugates; and characterization of DNA origami structures; TEM images of peptide-mediated DNA assemblies and determination of Kd values; error analysis (PDF)

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Notes

The authors declare no competing financial interest.

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